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## (54) Piperidine derivatives

(57) Novel compounds of formula

high blood pressure or as anti-depressants.

or pharmaceutically acceptable salts thereof, wherein Ar represents an optionally substituted indolyl group; A represents a straight or branched chain alkylene or oxoalkylene group, each having 2–4 carbon atoms and R represents an optionally substituted aryl (including heteroaryl) group, possess anti-hypertensive and psychotropic activity, and are useful in the treatment of

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## **SPECIFICATION**

## Pip ridin d rivatives

- 5 This invention relates to piperidine derivatives, to processes for preparing them and to pharmaceutical compositions containing them. This invention also relates to pyridinium and tetrahydro pyridinium compounds which are useful as intermediates in the preparation of the piperidine derivatives.
- More particularly this invention provides 4-acylaminopiperidyl derivatives which exhibit 10 pharmaceutical activity, especially antihypertensive activity in standard test procedures and also psychotropic activity as evidenced by their ability to inhibit parachloramphetamine induced hyperactivity. The piperidine compounds are therefore potentially useful in the treatment of high blood pressure or as antidepressants.

Accordingly this invention provides piperidine derivatives of formula

$$Ar-A-N$$
-NHCOCH<sub>2</sub>NHCOR (1) 20

- and pharmaceutically acceptable salts thereof, wherein Ar represents an optionally substituted indolyl group; A represents a straight or branched chain alkylene or oxoalkylene group each having 2 to 4 carbon atoms; and R represents an aryl (including heteroaryl) group which may be 25 substituted.
- Examples of groups for Ar are indol-3-yl which may be substituted by one or more groups selected from halogen, e.g. fluorine, chlorine or bromine (such as 5-chloro); alkyl having 1 to 6 carbon atoms, e.g. methyl, ethyl and propyl (such as 5-methyl); alkoxy having 1 to 6 carbon 30 atoms, nitro and hydroxy.
  - Examples of R are phenyl and phenyl substituted by the same groups as mentioned for the radical Ar. Heteroaryl R radicals include radicals where the heteroatom is nitrogen, such as pyridyl (e.g. pyrid-4-yl; sulphur, e.g. thien-2-yl; or oxygen, e.g. furan-2-yl. Heteroaryl R radicals may carry substituents as mentioned for the radical Ar.
- Examples of A are  $-(CH_2)_n$  where n is 2 or  $-CO(CH_2)_n$  where n is 1 to 3, e.g. oxobutylene. Pharmaceutically acceptable salts of the compounds of formula I include acid addition salts formed with inorganic or organic acids such as the sulphate, hydrochloride, hydrobromide, hydroiodide, nitrate, phosphate, sulphonate (such as the methanesulphonate or p-toluene-sulphonate), acetate, maleate, fumarate, tartrate and formate. Quaternary ammonium salts are 40 also included such as those formed with alkyl or aralkyl halides e.g. benzyl chloride, methyl iodide.
  - This invention also provides processes for preparing compounds of formula (I). In general the compounds of formula (I) can be made prepared by building up the molecule from appropriate starting materials in known manner.
- 45 One such process for preparing compounds of formula (I) as defined above comprises acylating a compound of formula:

55 wherein Ar and A are as hereinbefore defined, or a reactive derivative thereof, with an acid of formula:

HOOCCH₂NHCOR (IV)

60 wherein R is as hereinbefore defined, or a reactive derivative thereof. Coupling agents such as dicyclohexyl-carbodiimide may be used to effect acylation. As examples of the reactive derivatives of the acid of formula (IV) useful in the above m ntioned reaction mention is made of the acid halides, e.g. chloride, the azide and also 2-aryloxazol-5-ones of formula (IVa)

where R is as defined above.

Examples of the compound of formula (II) where the amino function is activated include the phosphazo derivative which may be coupled directly to the acid of formula (IV).

15 Compounds of formula (II) may be prepared according to processes described in UK Patent Specification Nos. 1218570 and 1345872.

A further process for preparing compounds of formula (I) as defined above comprises reacting a compound of general formula:

wherein Ar and A are as defined above and Y represents a leaving group, e.g. a halogen atom or an equivalent replaceable radical, e.g. a sulphonyloxy radical such as tosyloxy, with a compound of formula

$$+N$$
-NHCOCH<sub>2</sub>NHCOR  $(V$ 

wherein R is as hereinbefore defined. Further examples of Y when Ar is indol-3-yl and A is 35 -CH<sub>2</sub>- are disubstituted amino radicals such as dimethylamino or trisubstituted ammonium

radicals such as trimethylammonium (\*NMe<sub>3</sub>).

Yet a further process for preparing a compound of formula (I) comprises aroylating a

compound of formula

$$Ar - A - N$$
  $\longrightarrow$   $NHCOCH_2NH_2$ 

with an aroylating agent containing the group—COR wherein R is as hereinbefore defined e.g. using aroyl halides, aroyl anhydrides. Compounds of formula (VII) may be prepared by removing the α-amino protecting group from a corresponding compound of formula

$$_{55}$$
 Ar — A  $-$ N $\longrightarrow$ NHCOCH $_2$ NHB(VIII)

where Ar and n are as hereinbefore defined and B is an α-amino protecting group, e.g. benzyloxycarbonyl, t-butyloxycarbonyl. Methods for removing protecting groups and the protecting groups themselves are described in the standard textbooks on peptide chemistry, see for example e.g. E. Schroder and K. Lubke, "The Peptides" Volume I, Academic Press, New York and London, 1965. Compounds of formula (VIII) may be prepared by coupling a compound of

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formula (II) as hereinbefore defined with a compound of formula HOOCCH<sub>2</sub>NHB where B is as hereinbefore defined.

Compounds of formula (I) may also be prepared by treating a corresponding compound of formula (IX)

$$A_{r} \xrightarrow{A} \xrightarrow{\bigoplus} N \xrightarrow{N} -NHCOCH_{2}NHCOR$$

$$R^{l} \xrightarrow{D} \Theta \qquad (TX)$$

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to remove R<sub>1</sub>, wherein Ar and R are as hereinbefore defined, B⊖ is an anion as hereinbefore defined and R¹ is an organic quaternizing group which can be removed under mild conditions, e.g. by hydrogenolysis, that do not affect the rest of the molecule. For example, when R¹ is an arylmethyl radical, such as benzyl, hydrogenolysis under standard conditions, e.g. using an appropriate catalyst such as a palladium on carbon, platinum or nickel catalyst, gives compounds of formula (I). Methods for effecting this reaction are given in our U.K. Patent Specification No. 1,399,608. Suitable solvents include alkanols such as methanol.

Starting materials of formula (IX) maybe prepared by reacting a compound of formula (V) as defined above with a compound of formula

35 wherein R and R¹ are as defined above, with heating.
Compounds of formula I may also be prepared by reducing a corresponding compound of

formula (XI) or (XII):

$$A_r - A - N - NHCOCH_2NHCOR$$

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in which formulae Ar, A and R are as hereinbefore defined and B<sup>-</sup> represents an anion, e.g. a halide ion. For example catalytic hydrogenation e.g. in the presence of Raney nickel or platinum catalyst gives piperidine compounds of formula (I). The reduction may also be effected by a process described and claim d in our U.K. Patent Specification No. 1542137. Such a reduction process employes an alkali metal borohydride in a secondary alkanol having 3–5 carbon atoms, e.g. isopropanol. Alternatively reduction of compounds of formula (XII) using an alkali metal borohydride in methanol gives compounds of formula (XI).

Compounds of formula (XI) and (XII) are also within the scope of this invention. They may be 65 prepared by reacting compounds of formula (XIII) and (XIV)

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Sterile liquid form compositions include sterile solutions, suspensions, emulsions, syrups and elixirs. The active ingredient can be dissolved or supended in a pharmaceutically acceptable 65 sterile liquid carrier, such as sterile water, sterile organic solvent or a mixture of both. Preferably

a liquid carrier is one suitable for parenteral injection. Where the active ingredient is sufficiently soluble it can b dissolved in normal saline as a carrier; if it is too insoluble for this it can often be dissolved in a suitable organic solvent, for instance aqueous propylene glycol or polyethylene glycol solutions. Aqueous propylene glycol containing from 10 to 75% of the glycol by weight 5 is generally suitable. In other instances compositions can be made by dispersing the finely-5 divided active ingredient in aqu ous starch or sodium carboxylmethyl cellulose solution, or in a suitable oil, for instance arachis oil. Liquid pharmaceutical compositions which are sterile solutions or suspensions can be utilised by intramuscular, intraperitoneal or subcutaneous injection. In many instances, a compound is orally active and can be administered orally either 10 in liquid or solid composition form. 10 Preferably the pharmaceutical composition is in the unit dosage form. In such form, the composition is subdivided in unit doses containing appropriate quantities of the active ingredients; the unit dosage form can be packaged composition, the package containing specific quantities of composition, for example packeted powders or vials or ampoules. The unit dosage 15 form can be a capsule, cachet or tablet itself, or it can be the appropriate number of any of 15 these in package form. The quantity of active ingredient in a unit dose of composition may be varied or adjusted from 5 mg or less to 500 or more, according to the particular need and the activity of the active ingredient. The invention also includes the compounds in the absence of carrier where the compounds are in unit dosage form. A further aspect of this invention provides chemical intermediates for the compounds of 20 20 formula (I) which intermediates have the formula (XI) and (XII) as hereinbefore defined. The following Examples illustrate the invention: EXAMPLE 1 25 N-[2-[[1-[2-(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]benzamide 25 4-Amino-1-(2-[indol-3-yl]ethyl)piperidine (1.21g, 5mmol) and 2-phenyloxazol-5-one (0.8g, 5mmol; prepared according to the method of Stewart and Wooley, J.A.C.S. 78 5336 1956) were refluxed in methyl cyanide (30 cm<sup>3</sup>). After 1\(\frac{1}{2}\) hours more 2-phenyloxazol-5-one (0.1 g, 0.62 mmol) in methyl cyanide (10 cm<sup>3</sup>) was added. A further portion of 2-phenyl-oxozol-5-one 30 30 (0.1 g, 0.62 mmol) was added after 1 hour, and refluxing was continued for 30 minutes. The mixture was filtered hot and the filtrate on cooling gave the crude title compound which was collected and dried (1.55 g). The solid obtained was refluxed in isopropyl alcohol containing ethanolic HCl for three quarters of an hour and to give the hydrochloride salt which was collected after cooling overnight. This was sucked dry on the sinter, triturated with remuxing 35 ethanol for half an hour, collected, then triturated with ethanol containing 5-10% water, filtered 35 hot and dried to give the hydrochloride salt of the title compound (0.77g) mp 236-240°C. Analysis Found C, 64.50; H, 6.54; N 12.57%  $C_{24}H_{28}N_4O_2$  HCl,  $\frac{1}{4}H_2O$  requires: C, 64.71; H, 6.68; N, 12.58%. 40 40 **EXAMPLE 2** N-[2-[[1-[2-(1H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]-4-pyridinecarboxamide (i) Chloroacetyl chloride (4.0 cm<sup>3</sup>, 50.18 mmol) was added dropwise to a vigorously stirred mixture of 4-amino-1-(2-[indol-3-yl]ethyl)piperidine (12.10g, 49.8 mmol), potassium carbonate 45 (7.0g 50·72 mmol) water (100 cm³) and dichloromethane (300 cm³). After  $\frac{3}{4}$  hour more 45 chloroacetyl chloride (0.5 cm<sup>3</sup>, 6.27 mmol) and potassium carbonate (0.5g, 3.62 mmol) were added and stirring continued for a further 3 hour. The organic phase was separated, washed with water, dried over magnesium sulphate and evaporated to give 2-chloro-N-[1-[2-(1 H-indol-3yl)-ethyl]-4-piperidinyl]acetamide (19.29g crude) 50 (ii) A solution of the chloro product (3.5 g, 10.95 mmol) in strong ethanolic ammonia (140 cm³) 50 was heated at 100°C in a bomb for 22h. This was evaporated to give a glass (3.23g) which was purified by chromatography on silica (100-200 aktiv) eluting with chloroform 83: methanol 15: triethylamine 2 parts. This gave 0.92g of 2-amino-N-[1-[2-(1H-indol-3-yl)ethyl]-4-piperidinyl]acetamide. 55 (iii) The product of step (ii) is acylated with 4-pyridoyl chloride, hydrochloride to give the title 55 compound. m.p. 211-215°C. **EXAMPLE 3** The following procedure was used to test compounds of formula I for their ability to inhibit p-60 chloroamphetamine induced hyperactivity. 60 Three groups of 4 femal mic (20-24 g) received the test compounds (50 mg/kg po) and a

fourth group the requisite volume of vehicle. Thirty minutes later all the animals are given 20 mg/kg p-chloroamphetamine (pCA) ip. The grouped mice are placed immediately in square plastic cages in activity monitors and their motor activity recorded over the period 10-30 minutes post pCA. This procedure is repeated three more times so that four groups of mice are

may be substituted.

pyrid-4-yl group, each of which may be substituted.

used per treatment and each activity monitor is used with all treatments in turn. The inhibition of pCA induced hyperactivity is calculated thus:-

5 C-T 100% 5 C where C = mean activity of control groups 10-30 minutes post pCA. T = mean activity of treated groups 10-30 minutes post pCA. 10 This test is used as an in vivo screen for detection of 5-hydroxytryptamine uptake inhibitors. Compounds giving > 50% inhibition are considered of special interest. In such a test the compound of Example 1 showed 51.6% inhibition at 50 mpk. 15 EXAMPLE 4 15 The following procedure was used to test compounds of formila I for antihypertensive activity. Female rats are rendered hypertensive by implanting subcutaneously two wax pellets (30 mg) containing desoxycorticosterone acetate (15 mg) followed immediately by uninephrectomy. The drinking water is replaced by normal saline ad lib for 4 weeks. Blood pressures stabilise at a 20 hypertensive level after 6 weeks. Systolic pressure is measured directly before dosing with a test 20 compound using an E and M pneumatic pulse transducer and a Devices MX2 recorder. Groups of 4 rats are dosed orally with suspensions or solutions of test compound in 0.5% hydroxypropylmethylcellulose 0.9% saline vehicle. Blood pressures are recorded again at various time intervals and the results, expressed as a percentage of the pre-dose values compared with those 25 of a similar group of rats receiving vehicle alone. 25 In the above test the compound of Example 1 at a dose level of 50 mpk gave a 36.4% decrease in blood pressure after 2 and 6 hours. In the same test heart rate was decreased by 35.9% and 37.7% and 6 hours respectively after dosing. 30 EXAMPLE 5 30 Using a procedure analogous to Example 2, the following compounds may be reacted with 2amino-N-[1-[2-(1 H-indol-3-yl)ethyl]-4-piperidinyl]-acetamide benzoyl chloride 35 b) 4-chlorobenzoyl chloride 35 4-methoxybenzoyl chloride c) d) 2-thenoyl chloride 3-methylbenzoyl chloride e) to give: N-[2-[[1-[2(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]benzamide 40 a) 40 N-[2-[[1-[2-(1 H-indol-3-yl)ethyl-4-piperidinyl]amino]-2-oxoethyl]-4-chlorobenzamide. b) N-[2-[[1-[2(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]-4-methoxybenzamide. c) N-[2-[[1-[2(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]-2-thiophenecarboxamide. N-[2-[[1-[2(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]-3-methylbenzamide. e) 45 **CLAIMS** 1. A compound of formula 50 50 ([])55 55 or a pharmaceuticlly acceptable salt thereof wherein Ar represents an optionally substituted indolyl group; A represents a straight or branched chain alkylene or oxoalkylene group, each having 2 to 4 carbon atoms; and R represents an optionally substituted aryl or heteroaryl group. 2. A compound as claimed in Claim 1 wherein R represents -CH2CH2- or -COCH2CH2CH2-3. A compound as claimed in Claim 1 or Claim 2 wherein Ar is an indol-3-yl group which 60

4. A compound as claimed in any one of claims 1 to 3 wherein R represents a phenyl or

65 more groups selected from halogen, alkyl of 1 to 6 carbon atoms, alkoxy of 1 to 6 carbon

5. A compound as claimed in any one of claims 1 to 4 in which the substituents ar one or

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atoms, trifluoroalkyl, nitro and hydroxy.

- 6. N-[2-[[1-[2-(1 H-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl] benzamide or a pharmaceutically acceptable salt thereof.
- 7. N-[2-[[1-[2-(1 *H*-indol-3-yl)ethyl]-4-piperidinyl]amino]-2-oxoethyl]-4-pyridinecarboxamide or 5 a pharmaceutically acceptable salt thereof.
  - 8. A compound as claimed in any one of claims 1 to 7 which is in the form of a salt selected from sulphate, hydrochloride, hydrobromide, hydroiodide, nitrate, phosphate, methane-sulphonate, p-toluene-sulphonate, acetate, maleate, fumarate, tartarate and formate.
- 9. A process for preparing a compound of formula I as defined in Claim 1 which comprises:

  10 a) acylating a compound of formula

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$$Ar - A - N - NH_2$$
(II)

- wherein Ar and A are as defined in Claim 1 or a reactive derivative thereof, with an acid of 20 formula:
  - HOOCCH₂NHCOR (III)
- wherein R is as defined in Claim 1, or a reactive derivative thereof, or 25 b) reacting a compound of formula 25
  - Ar—A—Y (V)
- wherein Ar and A are as defined above and Y represents a leaving group, with a compound of 30 formula 30
- 35 HN NHCOCH<sub>2</sub>NHCOR (VI)
- 40 wherein R is as defined in Claim 1, or
  c) aroylating a compound of formula

- 50 with an aroylating agent containing the group -COR wherein R is as defined in Claim 1; d) treating a corresponding compound of formula IX
- 65 to remove R¹, wherein Ar and R are as defined in Claim 1, B<sup>⊙</sup> is an anion and R¹ is an organic

quaternizing group which can be removed under mild conditions that do not affect the rest of the molecule; or

e) reducing a corresponding compound of formula XI or XII

$$Ar - A - N - NHCOCH_2NHCOR$$
10 (XI)

in which formulae Ar, A and R are as defined in Claim 1 and B= represents an anion; or f) reacting a compound of formula:

wherein Ar and A are as defined in Claim 1 with a compound of formula VI is defined hereinabove in the presence of a catalyst; or

- 30 g) converting a base of formula I to a pharmaceutically acceptable salt thereof or *vice versa*.

  10. A compound of formula I whenever prepared by a process as claimed in Claim 9.
  - 11. A compound of formula I substantially as hereinbefore described in either Example 1 or Example 2(ii).
- 12. A compound of formula I as claimed in any one of claims 1 to 8 for use as an anti35 hypertensive agent.
  13. A compound of formula I as claimed in any one of claims 1 to 8 for use as an anti-
- depressant.

  14. A pharmaceutical composition comprising a compound of formula I as defined in any one of claims 1 to 8 or a pharmaceutically acceptable salt thereof and a pharmaceutical

  40 acceptable carrier.

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